

(FILE 'HOME' ENTERED AT 11:40:23 ON 16 APR 2003)

FILE 'MEDLINE, CAPLUS, BIOSIS, AGRICOLA' ENTERED AT 11:40:27 ON 16 APR 2003

L1 202587 S METHANOL  
L2 48 S L1 AND CULTURE AND INACTIVE  
L3 25 DUP REM L2 (23 DUPLICATES REMOVED)

FILE 'STNGUIDE' ENTERED AT 11:43:45 ON 16 APR 2003

FILE 'MEDLINE, CAPLUS, BIOSIS, AGRICOLA' ENTERED AT 12:39:17 ON 16 APR 2003

L4 3515 S GALACTOSE (2N) OXIDASE  
L5 78 S L4 AND (OXIDIZING)  
L6 8 S L5 AND CATALYTIC  
L7 5 DUP REM L6 (3 DUPLICATES REMOVED)

FILE 'STNGUIDE' ENTERED AT 12:40:22 ON 16 APR 2003

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FILE 'MEDLINE, CAPLUS, BIOSIS, AGRICOLA' ENTERED AT 11:40:27 ON 16 APR 2003

L1 202587 S METHANOL  
L2 48 S L1 AND CULTURE AND INACTIVE  
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FILE 'STNGUIDE' ENTERED AT 12:40:22 ON 16 APR 2003

FILE 'MEDLINE, CAPLUS, BIOSIS, AGRICOLA' ENTERED AT 13:01:03 ON 16 APR 2003

L8 340 S METHANOL (3N) INDUCTION  
L9 8 S L8 (10N) TEMPERATURE  
L10 4 DUP REM L9 (4 DUPLICATES REMOVED)

FILE 'STNGUIDE' ENTERED AT 13:02:05 ON 16 APR 2003

FILE 'CAPLUS' ENTERED AT 13:03:23 ON 16 APR 2003

FILE 'MEDLINE, CAPLUS, BIOSIS, AGRICOLA' ENTERED AT 13:03:26 ON 16 APR 2003

L11 246 S METHANOL AND INDUCTION AND TEMPERATURE  
L12 17 S L11 AND PICHIA  
L13 10 DUP REM L12 (7 DUPLICATES REMOVED)  
L14 118 S L11 AND (25 OR DECREAS? OR LOW?)  
L15 97 DUP REM L14 (21 DUPLICATES REMOVED)  
L16 4 S L15 AND PICHIA  
L17 1478 S METHANOL AND INDUC? AND TEMPERATURE  
L18 31 S L17 AND PICHIA  
L19 14 S L18 AND (25 OR DECREA? OR LOW?)  
L20 8 DUP REM L19 (6 DUPLICATES REMOVED)

AN 1993:18335 CAPLUS

DN 118:18335

TI Preparation of fully oxidized active and reduced inactive forms of **galactose oxidase** from *Dactylium dendroides* using ferricyanide-containing **oxidizing** and ferrocyanide-containing reducing forms of ion exchange resins

AU Montague-Smith, Michael P.; Wachter, Rebekka M.; Branchaud, Bruce P.

CS Dep. Chem., Univ. Oregon, Eugene, OR, 97403, USA

SO Analytical Biochemistry (1992), 207(2), 353-5

CODEN: ANBCA2; ISSN: 0003-2697

DT Journal

LA English

AB **Galactose oxidase** (EC 1.1.3.9) is a type II mononuclear copper protein secreted by the fungus *Dactylium dendroides*. The enzyme catalyzes the oxidn. of primary alcs. with O<sub>2</sub>, producing aldehydes and H<sub>2</sub>O<sub>2</sub>. The details of the **catalytic** mechanism have not been fully elucidated. A chronic problem in kinetic assays of **galactose oxidase** is the tendency of the enzyme to exist as a mixt. of oxidized active and one-electron reduced inactive forms. The two forms cannot be phys. sepd. by std. purifn. techniques. The enzyme can be activated by treatment with one-electron oxidants. This treatment results in maximally active enzyme. Once the active enzyme has been formed, the **oxidizing** agent must be removed, since it can substitute for the natural oxidant, dioxygen, in the enzyme reaction and alter reaction rates. Dialysis is tedious and does not lead to consistently reproducible results. Desalting columns are more effective but are not sufficiently consistent. Finally, activated enzyme is slowly reduced to a mixt. of active and inactive forms, so that a large quantity of identical enzyme necessary for extensive kinetic analyses cannot be maintained. A method of rapidly activating or deactivating small samples of enzyme to a consistent specific activity was developed. Ferricyanide and ferrocyanide, highly charged redox active anions, bind strongly to anion exchange resins, producing redox-active resins that are capable of **oxidizing** or reducing **galactose oxidase** to provide the active or inactive forms of the enzyme. The redox resins are easily made and stable and give reproducibly active or inactive enzyme.

L8 ANSWER 14 OF 18 CAPLUS COPYRIGHT 2003 ACS  
AN 1997:685007 CAPLUS  
DN 127:358067  
TI Online monitoring and control of **methanol** concentration in  
shake-flask **cultures** of *Pichia pastoris*  
AU Guarna, M. M.; Lesnicki, G. J.; Tam, B. M.; Robinson, J.; Radziminski, C.  
Z.; Hasenwinkle, D.; Boraston, A.; Jervis, E.; MacGillivray, R. T. A.;  
Turner, R. F. B.; Kilburn, D. G.  
CS Biotechnology Laboratory, University of British Columbia, Vancouver, BC,  
V6T 1Z3, Can.  
SO Biotechnology and Bioengineering (1997), 56(3), 279-286  
CODEN: BIBIAU; ISSN: 0006-3592  
PB Wiley  
DT Journal  
LA English  
AB The methylotrophic **yeast** *P. pastoris* can be used to express  
recombinant genes at high levels under the control of the MeOH-  
**inducible alc. oxidase 1 (AOX1) promoter**.  
Accurate regulation of the MeOH concn. in *P. pastoris* **cultures**  
is necessary to maintain induction, while preventing accumulation of MeOH  
to cytotoxic levels. An inexpensive MeOH sensor that uses a gas-permeable  
silicone rubber tube immersed in the **culture** medium and an org.  
solvent vapor detector was developed. The sensor was used to monitor MeOH  
concn. continuously throughout a fed-batch shake-flask **culture**  
of a *P. pastoris* clone producing the N-lobe of human transferrin. The  
sensor calibration was stable for the duration of the **culture**  
and the output signal accurately reflected the MeOH concn. detd. off-line  
by HPLC. A closed-loop control system utilizing this sensor was developed  
and used to maintain a 0.3% MeOH concn. in the **culture**. Use of  
this system resulted in a 5-fold increase in volumetric protein  
productivity over levels obtained using the conventional fed-batch  
protocol.

L15 ANSWER 1 OF 1 MEDLINE DUPLICATE 1  
 AN 96292476 MEDLINE  
 DN 96292476 PubMed ID: 8728322  
 TI Expression and secretion of rabbit plasma cholesteryl ester transfer protein by *Pichia pastoris*.  
 AU Kotake H; Li Q; Ohnishi T; Ko K W; Agellon L B; Yokoyama S  
 CS Lipid and Lipoprotein Research Group, University of Alberta, Edmonton, Canada.  
 SO JOURNAL OF LIPID RESEARCH, (1996 Mar) 37 (3) 599-605.  
 Journal code: 0376606. ISSN: 0022-2275.  
 CY United States  
 DT Journal; Article; (JOURNAL ARTICLE)  
 LA English  
 FS Priority Journals  
 EM 199611  
 ED Entered STN: 19961219  
 Last Updated on STN: 19990129  
 Entered Medline: 19961125  
 AB The rabbit cholesteryl ester transfer protein (CETP) was expressed in the methylotrophic yeast *Pichia pastoris* by introducing the CETP cDNA under the control of the **methanol-inducible** alcohol oxidase **promoter**. The cDNA was cloned from in vitro amplified cDNA of rabbit liver mRNA. The nucleotide sequence of the cloned cDNA differed slightly from the previously published sequence that changed the amino acid sequence in six residues. Interestingly, five of these replacements are identical to the corresponding residues in human CEPT. In addition, the encoded mature N-terminal sequence was changed from Cys- to Arg-Glu-Phe- to link the CETP sequence to the yeast acid phosphatase signal peptide. The culture medium of the transformed cells induced with 1% methanol contained both cholesteryl ester and triglyceride transfer activity comparable to that of rabbit plasma. Like rabbit plasma, the lipid transfer activity in the medium could be inhibited by monoclonal antibodies that block CE/TG transfer or TG transfer alone. Immunoblot analysis of M(r) = 80 K and minor species of M(r) = 60-100 K. In spite of these differences, the specific transfer activity of the recombinant CETP was indistinguishable from that of rabbit plasma CETP of M(r) = 74 K. N-Glycosidase F treatment converted both the recombinant and plasma CETP to a single species of M(r) = 55 K. Both the plasma and recombinant CETP lost their activity after removal of N-linked carbohydrate and sialic acid. A single 55 K component was found in the cell-lysates. The intracellular form of the recombinant CETP was not modified by N-glycosidase F treatment. In conclusion, the recombinant CETP is synthesized as an **inactive** polypeptide that is processed and secreted as a functional glycoprotein. In addition, the N-terminal Cys residue of the plasma CETP is not required for its activity.